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# High-throughput retrotransposon-based genetic diversity of maize germplasm assessment and analysis

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## Abstract

Maize is one of the world's most important crops and a model for grass genome research. Long terminal repeat (LTR) retrotransposons comprise most of the maize genome; their ability to produce new copies makes them efficient high-throughput genetic markers. Inter-retrotransposon-amplified polymorphisms (IRAPs) were used to study the genetic diversity of maize germplasm. Five LTR retrotransposons (*Huck*, *Tekay*, *Opie*, *Ji*, and *Grande*) were chosen, based on their large number of copies in the maize genome, whereas polymerase chain reaction primers were designed based on consensus LTR sequences. The LTR primers showed high quality and reproducible DNA fingerprints, with a total of 677 bands including 392 polymorphic bands showing 58% polymorphism between maize hybrid lines. These markers were used to identify genetic similarities among all lines of maize. Analysis of genetic similarity was carried out based on polymorphic amplicon profiles and genetic similarity phylogeny analysis. This diversity was expected to display ecogeographical patterns of variation and local adaptation. The clustering method showed that the varieties were grouped into three clusters differing in ecogeographical origin. Each of these clusters comprised divergent hybrids with convergent characters. The clusters reflected the differences among maize hybrids and were in accordance with their pedigree. The IRAP technique is an efficient high-throughput genetic marker-generating method.

**Keywords** *Zea mays* · Biodiversity · Transposable elements · Retrotransposons

**Electronic supplementary material** The online version of this article (<https://doi.org/10.1007/s11033-020-05246-4>) contains supplementary material, which is available to authorized users.

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## Abbreviations

IRAP	Inter-retrotransposon-amplified polymorphism
TE	Transposable element
LTR	Long terminal repeat
RLX	LTR retrotransposon

## Introduction

The long terminal repeat (LTR) retrotransposons (RLX) [1, 2] are a large class of transposable elements that propagate in the genome by a “copy-and-paste” mechanism that is essentially identical to the intracellular phase of retrovirus replication [1–5], in contrast to the “cut-and-paste” mobility of DNA transposons. The RLX lifecycle involves transcription of an integrated copy, reverse transcription of the transcript into cDNA, and integration of the new copy. Because the RLX mother copy remains part of the chromosome and the daughter copies integrate at new loci, the precise insertion points for the daughter are unlikely to be identical in lines diverging by descent. Complete understanding of the genome and the relationship between genotype and phenotype requires knowledge of both the role and function of the genes as well as of the repetitive component, particularly regarding RLX dynamics [3]. Most eukaryotic genomes comprise over 70% repetitive DNA, with gene numbers, ranging from 10,000 to 50,000, showing much less variation at the monoploid level [4–7]. Particularly in higher plants, RLXs compose more than half of the repetitive DNA; they not only facilitate homologous recombination, but also can undergo intra- and inter-RLX recombination that is part of their dynamism [4, 8–10]. The RLXs are generally dispersed throughout genome, displaying relatively high structural diversity [11–15]. Retroelements have been suggested as an important creative force in genome evolution, driving processes such as mutation, recombination, genome expansion, and adaptation of an organism to changing environmental conditions [3, 14, 16].

All these properties make RLXs excellent sources of efficient high-throughput genetic markers in eukaryotic, especially plant, genomes [17]. The presence or absence of RLX insertions at particular loci can be surveyed by polymerase chain reaction (PCR)-based genotyping methods [17–20]. The RLX-based genetic marker techniques rely on PCR amplification between features such as the LTR that are conserved in RLXs, or between these features and other dispersed and conserved motifs in the genome. The methods include retrotransposon-based insertion polymorphism (RBIP) [21], sequence-specific amplified polymorphism (S-SAP) [22], inter-retrotransposon-amplified polymorphism (IRAP) [19],

retrotransposon-microsatellite-amplified polymorphism (REMAP), and inter-primer binding site (iPBS) amplification [17–19, 23–27].

These retrotransposon-based high-throughput genetic DNA fingerprinting methods are both highly informative and polymorphic, even in areas of chromosomes showing low levels of inter-genic recombination and therefore haplotypes with few genic single-nucleotide polymorphisms (SNPs); RLX markers are consistent with geographical and morphological data. The stability of retrotransposon integration sites and recombination events allows them to be used as molecular genetic markers in genetic map construction [28–32]. Retrotransposon markers have also been widely used to assess genetic diversity in many species [33–39]. Given that plant retrotransposons are stress-activated [15, 40], their role in generating ecogeographical patterns of genomic diversity is of particular interest. Retrotransposon markers have been applied successfully to the analysis of genetic diversity in various genera and species, such as apple, rice, sunflower, grapevine, flax, and alfalfa [38, 41, 42].

Maize (*Zea mays* L.) is a very special species among the cereal crops, because of its high phenotypic and genomic diversity [43]. Maize is important worldwide as a food and feed crop, and also as an energy crop, due to its high biomass potential. Moreover, it has long been used as a model organism for plant biology. Maize was the first eukaryote in which transposable elements (TEs) were discovered, during the mid-twentieth century, by the Barbara McClintock, bringing her the Nobel Prize [44]. TEs and, particularly, retrotransposons, comprise most of the maize genome; 95% of maize TEs are RLXs [43, 45, 46]. Through their copy number variation, rearrangements, and polymorphic loci, the TEs contribute most of the genome variation between maize lines, where amplification of a few retrotransposon families is the major cause of “genomic obesity” [47, 48]. High-throughput sequencing methodologies have demonstrated that some families of TEs show considerable transcriptional activity.

In the present study, we developed and applied a high-throughput IRAP technique for five RLX families to detect genetic polymorphisms among maize germplasm. These families (*Opie*, *Ji*, *Cinful*, *Huck*, and *Grande*) comprise a large fraction of the maize genome [49–54], up to 25% of the total. The main goal was to find efficient and high-throughput retrotransposon markers for diversity analyses and to assess the polymorphism of these markers among maize genotypes originating from different ecogeographical origins. The IRAP genetic markers developed were used to compare the genetic variability among maize cultivars and breeding lines differing in ecogeographical origin to detect correlations between phenotypic characters and retrotransposon markers.

## Materials and methods

### Plant material and DNA extraction

Grains of maize lines and hybrids were kindly provided by the Maize Research Section, Agricultural Research Center (ARC) and U.S. Department of Agriculture (USDA). The names of these hybrids are listed in Table 1 and Supplemental Data 1. Further data on the genotypes can be found on the National Germplasm Resources Laboratory homepage (<https://npgsweb.ars-grin.gov/gringlobal/taxonomygenus.aspx?id=13020>).

DNA was isolated from leaves of 10-day-old seedlings of the 16 hybrids using the cetyltrimethylammonium bromide (CTAB) extraction protocol with modifications (CTAB solution: 2% CTAB, 1.5 M sodium chloride (NaCl), 20 mM ethylenediaminetetraacetic acid trisodium salt (Na<sub>3</sub>EDTA), and 0.1 M (*N*-[2-hydroxyethyl]piperazine-*N'*-[2-ethanesulfonic acid]) (HEPES), pH 5.3), as described (<https://primerdigital.com/dna.html>), with RNase A treatment. A detailed protocol for DNA isolation was deposited at protocols.io [55]. The DNA samples were diluted in 1×TE buffer. The DNA quality was checked electrophoretically, as well as

spectrophotometrically with a Nanodrop apparatus (Thermo Fisher Scientific Inc., Waltham, MA, USA).

### TE sequence sources and PCR primer design

Thirty LTR primers were designed, based on the most abundant RLX groups in maize (Cinful1 (AC231746), *Huck1* (AC230001), *Ji* (DQ002406), *Opie* (AY664413), *Grande* (AY664416.1:70909-83340), and *Tekay* (AF050455)). The RLX sequences were obtained from the TRansposable Elements Platform (TREP) database (<https://botserv2.uzh.ch/kelldata/trep-db/>), and analysis of homologous sequences was performed on the output of the National Center for Biotechnology Information (NCBI) search results. For a given family of retrotransposons, their LTRs showed sequence variability, but certain regions were relatively conserved. For each family, the sequence accessions were aligned and conservation assessed with the multiple alignment procedure of MULTALIN [56]. The conserved segments of the LTR of the retrotransposons were used for the design of PCR primers, which was carried out with the program FastPCR (<https://primerdigital.com/fastpcr.html>) [57–59]. Several inverted primers at both ends of the LTRs of each retrotransposon to compare the efficiency and reproducibility of amplification were

**Table 1** List of the tested 16 maize hybrids and their commercial names

ID	Name	Pedigree	Kernel color
H-1	Sc122	Gz628 x Gz603	White
H-2	Sc124	Gz629 x Gz603	White
H-3	Sc10	Sd7 x Sd63	White
H-4	Sc168	Gz639 x Gz658	Yellow
H-5	Sc164	Gz639 x Gz655	Yellow
H-6	Sc162	Gz639 x Gz653	Yellow
H-7	Sc128	SK9 x SK5	White
H-8	Sc129	Gz612 x Gz628	White
H-9	Sc166	Gz639 x Gz656	White
H-10	TC 310	SC10 x Sd34 [(Sd7 x Sd63) x Sd43]	White
H-11	TC 311	S C21 x Sd34 [(Gm2 x Sd.63) x Sd34]	White
H-12	TC 314	SC24 x Sd34 [(Gm18 x Sd.63) x Sd34]	White
H-13	TC 321	SC21 x Sd7	White
H-14	TC 324	SC24 x Sd7	White
H-15	TC 323	SC23 x Sd7[(Gm4 x Sd63) x Sd7]	White
H-16	TC 352	SC52 x Gm1021 [(Gm1002x Gm1004) x Gm1021]	Yellow

SC single cross, TC triple cross, Gz Giza, Gm Gemmeza, Sd Sids, H hybrid (just cod letter)

Name: SC and TC = (name of hybrid in our institute)

Giza, Gemmeza and Sids = places of original hybrids

For example, = Gz603 = Egyptian maize hybrid produced in Giza

Pedigree: each cross like sc or TC produced from hybridization between two parents

SC122 = produced from hybridization between Gz628 and Gz603

TC310 = produced from hybridization between SC10 (single cross) and Sd34 (which produced from hybridization Sd7 and Sd63 multiple Sd43)

**Table 2** Primers used for IRAP

ID	Sequence	Location at LTR	bp	Tm (°C) <sup>a</sup>	CG (%)	LC (%)	PQ (%)
Huck (gypsy), LTR (1682 bp)							
6301	ggtctcgaacgccgaca	1665 → 1682	18	57.5	61.1	83	67
6302	ccctactgcaagcgacc	1488 → 1505	18	58.9	66.7	81	80
6303	gcgtggacgcttgaacc	1470 → 1487	18	56.9	61.1	89	89
6304	atgggggttaccagctgagaattagg	36 ← 61	26	60.1	50.0	89	78
6305	cccatcaggccttgcagc	71 ← 89	19	59.3	63.2	81	71
6306	tggccgaggctaggctcg	141 ← 158	18	61.2	72.2	72	62
4303	gacgtcctcttgcctcgtaagg	1149 ← 1171	23	60.0	56.5	88	78
4304	tgtcgagtccttgcctgacgt	1167 ← 1186	20	58.1	55.0	84	84
4305	agaggtttggcaccctgttcg	1246 ← 1267	22	59.9	54.5	88	88
4306	attactcgctcggtccaggac	1639 → 1659	21	58.1	57.1	87	77
4307	acgcctataaaaggaaggtcca	1368 → 1389	22	55.5	45.5	88	75
Ji (copia), LTR (1208 bp), DQ002406							
4308	cactcggcttaagtaccgt	246 ← 265	20	56.1	55.0	92	67
4309	gacccggctcttggaccac	201 ← 220	20	58.3	60.0	82	80
4310	ttgtgaccacctaaccgggag	189 ← 210	22	61.0	59.1	88	77
4311	ttgcaagaaccgaacctcgta	162 ← 183	22	58.5	50.0	90	67
4312	tgtgttgggcaattcaacca	1178 → 1197	20	55.1	45.0	82	80
Opie (copia), LTR (1293 bp), AY664413							
4313	caagtaccactaatcactggagc	628 ← 650	23	55.6	47.8	90	90
4314	tcccttggcgcctcataacc	608 ← 627	20	59.2	60.0	92	92
4315	tcataaccgcgagcagcgctgat	593 ← 615	23	62.9	56.5	95	88
4316	atcctgtttggctgtagtggac	854 ← 875	22	56.9	50.0	85	75
4317	gttcgcattcggttgcctt	1256 → 1275	20	57.1	50.0	79	79
4318	gtttgccttctaaaaccgggt	1267 → 1287	21	55.8	47.6	87	80
4319	tgatgtgtactcacccttgc	891 → 910	20	54.5	50.0	87	80
Grande (gypsy), LTR (644 bp), AY664416.1:70909-83340							
4320	ggtgtgcggctcgaccaca	604 → 622	19	62.2	68.4	78	73
4321	aagcatactcaccaggacgt	452 → 471	20	55.4	50.0	95	87
4322	gcattcttaagcgcccgaaactg	492 → 515	24	64.5	62.5	93	82
4323	ctaggccttatatactcgaccgt	328 ← 350	23	55.3	47.8	88	83
4324	ggctccagcgcggaagtc	197 ← 216	20	63.7	70.0	84	70
Tekay (gypsy), LTR (3441 bp), AF050455							
4325	gtgggtaaagttgtaccacctc	2768 → 2789	22	55.5	50.0	85	85
4326	ccctggaaccgggtgtgaca	3422 → 3441	20	61.1	65.0	82	70
4327	gtggaatggatcctggcac	3376 → 3395	20	56.2	55.0	82	67
4328	cagagccatgatagcactcgtg	2470 ← 2492	23	60.1	56.5	93	88
4329	ttatcctgggtgtcactccatgtt	2436 ← 2461	26	60.3	50.0	82	82
4330	ttaccacaagccatgtatcc	2755 ← 2775	21	54.5	47.6	87	87
Cinful1 (gypsy), LTR (628 bp), AC231746							
4331	gcatttgagaacaagtcacca	185 ← 212	21	55.5	47.6	87	73
6307	ctaaaaggctattctatcttaagtcggtt	132 ← 161	30	55.8	33.3	75	75
4332	acaattatgccctcataaaagttaca	605 → 625	28	55.6	32.1	85	67

<sup>a</sup>Tm—the melting temperature, calculated with 0.4 μM concentration of primer and without Mg<sup>2+</sup>

designed. The sequences of the primers are shown in Table 2. None of the primers chosen formed self-dimers, and all showed high PCR efficiency for IRAP fingerprinting. The chosen primers matched the motifs sufficiently

conserved in the retrotransposons to allow amplification of the great majority of targets in the genome.

## PCR analysis

IRAP analysis was conducted according to Kalendar and Schulman [19], using 30 primers from the LTRs of the five RLXs. The PCRs were performed in 25- $\mu$ L reaction mixtures containing 25 ng genomic DNA, 1  $\times$  DreamTaq buffer, 200 mM deoxyribonucleotide triphosphate (dNTP), 400 nM primer and 1 U DreamTaq DNA Polymerase (Thermo Fisher Scientific). The amplifications were performed in the MasterCycler Gradient (Eppendorf AG, Hamburg, Germany). The PCR reaction program consisted of 1 cycle at 95 °C, 2 min; 30 cycles of 95 °C for 15 s, 60 °C for 20 s, 72 °C for 40 s; and a final elongation at 72 °C for 2 min.

The PCR products were separated by electrophoresis at 70 V for 8–10 h in a 1.4% agarose gel (Wide Range; SERVA Electrophoresis GmbH, Heidelberg, Germany) with 1  $\times$  Tris-HEPES-EDTA (THE) electrophoresis buffer [18]. The Thermo Scientific GeneRuler DNA Ladder Mix, 100–10,000 base pairs (bp), #SM0332, was used as a standard. The gels were stained with ethidium bromide (EtBr) and scanned, using an FLA-5100 imaging system (FUJi Photo Film GmbH; now FUJIFILM Europe GmbH, Heidelberg, Germany) with a resolution of 50  $\mu$ m.

## Data analysis

From the IRAP fingerprint profiles, all clear bands were scored at each band position for each primer in all samples. Polymorphic bands (PBs) of the same size were assumed to represent a single locus. The presence or absence of a fragment of a given length was recorded in binary code. The gels were scored of a total of 677 PBs for the samples. Based on the primary data, the level of genetic diversity as defined by Nei was determined, using Arlequin software [60]. The method applied was based on cluster analysis expressing the relationships of the hybrids examined as a distance percentage in a cluster tree and similarity matrix. The data were analyzed, using NTSYSpC software (Numerical Taxonomy and Multivariate Analysis System) version 2.11 (<https://www.exetersoftware.com/cat/ntsyspc/ntsyspc.html>).

The primary genetic data were bootstrapped with SEQBOOT (<https://csbf.stanford.edu/phylip/seqboot.html>), after which the pairwise genetic distances were calculated, using the Genetic Distance Matrix Program (GENDIST) (<https://www.bablobk.de/gendist/>). Both programs are from PHYLIP (Phylogeny Inference Package) software package (<https://evolution.gs.washington.edu/phylip.html>). The ability of IRAP markers to reveal genetic relationships among all the maize accessions was evaluated phylogenetically by neighbor-joining (NJ), for which an algorithm was constructed, using PAUP software [61]. Support for the tree was determined by performing 1000 bootstrap operations on the dataset generated by distance analysis. To determine

the partitioning of the IRAP genetic variation into inter- and intrapopulation variance components, analysis of molecular variance (AMOVA) was conducted with the program Genetic Analysis in Excel (GenAlex) 6.5 [62].

Summary statistics related to the number of bands generated by each genotype (NTI) and for each group only, including the number of PBs, percentage of polymorphic loci (PPL%), number of private bands (NPB), Shannon's Information Index (I), genetic differentiation index (PhiPT) among populations, Nei's genetic distance (D), and Nei's genetic identity (IN), were calculated using GenAlex 6.5. Genetic distance, using minimum Jaccard coefficients, was calculated with Factor Analysis of Mixed Data (FAMD) 1.31. A dendrogram for the studied genotypes was constructed, based on the maximum likelihood method [63], using MEGA X software [64].

## Results and discussion

### In silico PCR analysis of the maize genome

We performed in silico IRAP analysis, using FastPCR software for the maize (B73 RefGen\_v4) (<https://www.ncbi.nlm.nih.gov/genome/12>) and sorghum (*Sorghum bicolor* (L.) Moench) (*Sorghum\_bicolor\_NCBIV3*) (<https://www.ncbi.nlm.nih.gov/genome/108>) genomes, using a single LTR primer corresponding to a sequence highly conserved in the RLXs examined. For in silico IRAP analysis, we applied the default options, since the length of potential PCR products varied from 50 to 3000 bp and allowed a single mismatch within the 3'-termini of the LTR primer. The results of the in silico IRAP analysis are represented in Table 3. As expected, no amplicons were predicted for the *Sorghum bicolor* genome, due to high divergence of the RLX sequences in the maize genome from those in *Sorghum bicolor*.

The number of in silico IRAP amplicons perfectly correlated with the results in the wet-lab IRAP experiments. The LTR primers that produced many in silico IRAP amplicons (over 50 per maize genome) also showed a strong background of multiple and overlapping IRAP amplicons in the experiments. For example, LTR primers 4309, 4310, 4311, and 4312 for the *Ji* RLX family produced the highest number of IRAP amplicons in the in silico analysis. As in the PCR experiments, we observed a strong background over the entire range of PCR fragments. For this reason, the IRAP amplicons with very high numbers in the PCR experiments could not be adequately resolved on an agarose gel, and these primers were not used further to analyze maize lines. LTR primer 4331 from Cinfu1 yielded over 389 IRAP amplicons in in silico analysis of the B73 line; in the PCR experiment, it also resulted in a fairly strong background over the entire range of PCR fragments. Similarly,



**Table 3** Total number of in silico IRAP hits returned by the search of maize genome (B73, RefGen\_v4) using LTR primer corresponding to a different LTR-retrotransposons

ID	Primer sequence	Chromosomes of B73 genome (RefGen_v4)										IRAP	
		ch1	ch2	ch3	ch4	ch5	ch6	ch7	ch8	ch9	ch10		Total bands
Huck (gypsy)													
6301	ggtctcgaaacgccgaca	14	17	12	8	14	16	5	7	11	5	109	
6302	ccctactgcaagcgacc	14	16	13	7	18	15	5	8	14	9	119	
6303	gcgtggacgcttgaacc	13	17	12	8	16	17	6	8	11	8	116	
6304	atgggggttaccagctgagaattagg	8	6	6	10	17	10	11	15	12	9	104	
6305	cccatcaggcctttgcagc	10	9	17	13	22	8	11	11	13	16	130	
6306	tggccgaggctaggctcg	9	6	5	7	16	7	9	13	11	11	94	
4303	gacgtcctcttgcttccgtaagg	5	6	5	2	3	4	6	9	9	10	59	56
4304	tgtcgagtccttgctgacgt	10	7	7	6	9	5	3	9	5	6	67	58
4305	agagggttttggcacccctgttcg	1	1	0	1	0	0	0	1	0	1	5	
4306	attactcgtcgggtccagggac	10	7	3	2	5	5	6	6	4	3	51	44
4307	acgcctataaaaggaaggtcca	5	6	6	0	1	0	1	4	3	3	29	23
Ji (copia)													
4308	cactcgggtctaagtaccgt	30	13	14	17	15	20	13	13	2	8	145	
4309	gacccgggtctttgtgaccac	51	30	22	31	32	34	21	23	11	21	276	49
4310	ttgtgaccacctcaacggggag	89	38	44	27	60	47	32	47	37	24	445	30
4311	ttgcaagaaccgaacctcggtta	103	69	54	52	67	57	42	58	39	39	580	32
4312	tgtgttgggcaattcaacca	93	45	42	43	58	37	41	49	37	30	475	38
Opie (copia)													
4313	caagtaccactaatcactggagc	0	0	1	1	0	0	0	1	1	1	5	
4314	tccttggccgctcataacc	1	1	2	3	2	0	0	3	2	5	19	
4315	tcataaccgcgagcacggctgat	4	3	2	4	3	2	0	0	4	3	25	30
4316	atcctgttttgctgtagtggac	1	0	2	3	3	2	3	2	2	3	21	
4317	gttcgattcggtttgccitt	10	14	12	15	12	11	4	4	6	4	92	45
4318	gtttgccttctaaaaccgggt	8	16	11	13	3	11	5	5	3	6	81	31
4319	tgatgtgtactacccttgc	14	27	19	29	16	14	13	12	12	11	167	20
Grande (gypsy)													
4320	ggtgtgcggtcggacccaa	4	2	0	1	3	0	2	0	0	0	12	29
4321	aagcatactaccaggacgt	4	1	0	1	1	0	1	0	0	1	9	
4322	gcattctctaagcgcccgaaacctg	5	3	1	1	2	0	4	1	1	3	21	23
4323	ctaggccttatatactgaccgt	4	1	0	0	1	0	2	3	0	2	13	21
4324	ggctccagcgcgggaaagtc	4	1	1	1	0	0	1	3	0	4	15	39
Tekay (gypsy)													
4325	gtgggttaaagttgtaccacctc	0	0	0	0	1	0	0	0	0	0	1	19
4326	ccctggaaccgggtgtgaca	16	27	17	20	16	15	9	11	9	10	150	24
4327	gtggaaatggatcctggcac	2	3	1	0	0	1	1	0	1	1	10	39
4328	cagagccatgatagcactcgtgg	1	0	0	0	0	0	1	0	1	0	3	27
4329	ttatcctgggtgtgctaccatgttc	3	2	0	1	1	1	2	1	2	0	13	
4330	ttaccacaagccatgtatcc	3	1	0	0	0	0	0	1	0	0	5	
Cinfull (gypsy)													
6307	ctaaaaggctattctatctttaagtcggtt	26	26	19	29	20	18	23	17	13	19	210	
4331	gcatttgagaacaagtcccca	46	49	46	51	48	25	34	32	24	34	389	
4332	acaattatgccctcataaaagtttaca	0	1	1	2	0	0	1	0	0	0	5	

LTR primer 4326 from *Tekay* produced over 150 IRAP amplicons in in silico analysis, while LTR primers 4317, 4318, and 4319 from the *Opie* RLX generated 92, 81, and 167 IRAP amplicons, respectively. Another important result from the in silico IRAP analysis is that it did not matter in which direction the primers were oriented at each terminus of the LTR; in either case, similar results were obtained. The number of IRAP amplicons was dependent only on the copy number of a particular RLX and how conserved the LTR region was for which the primers were designed.

### Diversity assay among Egyptian maize hybrids

The maize genome is composed of a diverse group of RLXs that are major sources of genetic variations. The selection of effective LTR primers must consider the abundance and distribution of the RLX family. In this study, the LTR primers used were designed for five different LTR retrotransposon families belonging to the high-copy classes *gypsy* and *copie*. LTR primers are usually designed to complement areas as close as possible to the 5' or 3' ends of the LTR, which is the most conservative part, and to minimize the amplification of long LTR fragments. The effectiveness of IRAP amplification was directly dependent not only on the total number of copies of the element, but also on the degree of LTR regional conservation. Screening for single primers resulted in the selection of 30 LTR primers for IRAP (Fig. 1). All selected LTR primers yielded 20–60 scorable bands and showed high-quality reproducible DNA fingerprints.

Overall, 677 amplicons were scored as effective, using all LTR primers from the five RLX families, of which 392 were polymorphic, with a mean polymorphism of 58%. A total of 40 unique IRAP amplicons were generated, including 22 positive and 18 negative unique bands, using 20 primers. Primers 4319 (*Opie*) and 4324 (*Grande*) produced the highest number of unique bands (7 and 6, respectively). The *Grande* RLX showed the highest number of unique bands (13), while the *Huck* LTR primers produced the smallest number (3). These results were expected and corresponded to the result of in silico IRAP analysis. The maize genome is relatively large (2182.61 Mb), and consists of about 50% RLX sequences. The size of the genome, with the main part comprising RLXs, increased the efficiency of the RLX-based methods and resulted in a high percentage of polymorphism, using a single primer. Due to the ability of retrotransposons to integrate into a multitude of loci in the genome, they constituted informative molecular markers for the plant species. The pattern obtained was related to the copy number and the size of the RLX family. The PCR products and the polymorphism resulted from the amplification of hundreds to many thousands of target sites in the genome. These polymorphisms functioned as means of identification, in detecting genetic erosion, and in revealing genetic relationships.

The LTR primers revealed different levels of polymorphism among the maize lines examined. Tables 4 and 5 showed that the LTR primers applied in 16 maize hybrids produced 677 bands; 392 of these were PBs and 285 were monomorphic bands. LTR primer 4304 (*Huck*) produced the highest number of bands (58) and primer 4325 (*Tekay*) the lowest number (19). The highest percentage of PBs was produced by primer 4310 (93%) for the *Ji* RLX and the lowest percentage by *Opie* LTR primer 4318 (26%). The number of polymorphic amplicons per primer ranged from 8 with 4318 (*Opie*) to 33 with 4306 (*Huck*). On average, the number of amplicons per primer throughout the 16 genotypes was 34 and for polymorphic amplicons were 20%. Various levels of polymorphism among primers were detected that ranged from 93% for primer 4310 (*Ji*) to 26% for primer 4318 (*Opie*) and were also recorded for each TE family. The primers based on the *Grande* elements showed the highest percentage of polymorphism (75%), compared with those based on the *Ji*, *Huck*, *Opie*, and *Tekay* elements, which showed total polymorphism of 55%, 53%, 48%, and 63%, respectively.

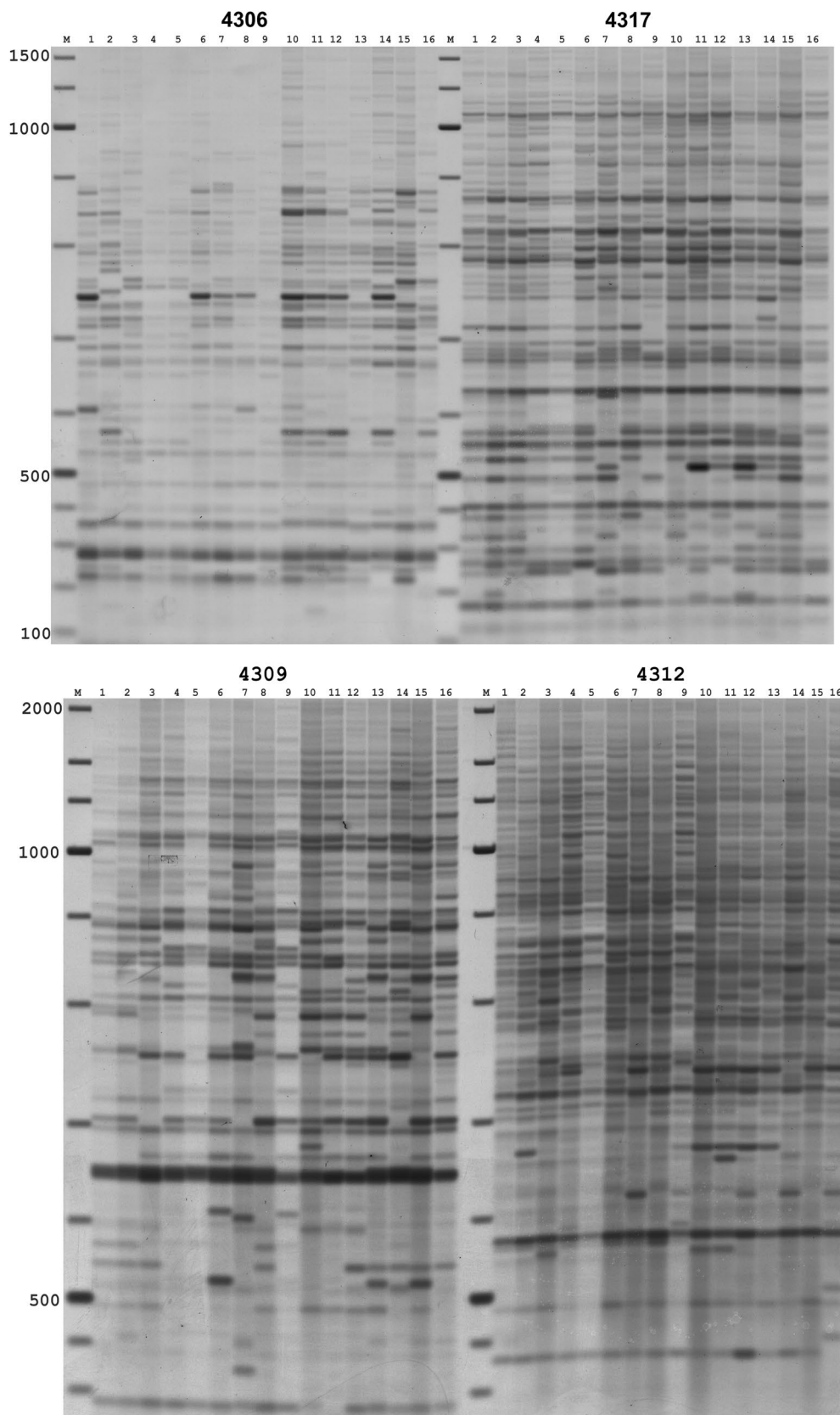
### IRAP markers based on the *Ji* family

Five LTR primers for the *Ji* RLX were tested in 16 maize hybrids, four of which (4309–4312) revealed variability among maize hybrids (Table 4). The bands ranged from 100 to 2000 bp. The total number of bands generated from the *Ji* element-based primers was 149; 82 were PBs with a mean polymorphism of 55%. Unique bands, either positive or negative (present or absent), characterized the maize hybrids. Five positive unique bands of 680, 2550, 1120, 410, and 500 bp characterized hybrids H-10, H-7, and H-16, whereas only one negative band of 298 bp, resulting from primer 4309, characterized H-11. Each band distinguished its respective hybrid and could be used as a fingerprint.

### IRAP markers based on the *Huck* family

Five LTR primers for the *Huck* RLX were tested, four of which (4303–4307) revealed variability among the maize hybrids (Table 4). A total of 181 number of bands were generated by the *Huck*-based primers, and ranged from 160 to 5100 bp; 96 PBs and 85 monomorphic bands were generated, with a mean of 53% polymorphic. The *Huck*-based LTR primers characterized H-15 and H-11 with two unique bands of 1301 and 1600 bp, respectively, while a single unique negative band of 2400 bp was generated from primer 4303 and characterized H-5 (Table 5). These bands could be used to fingerprint their respective genotypes.





**Fig. 1** Agarose gel electrophoresis patterns of IRAP amplicons obtained with primers 4306 (*Huck*), 4317 (*Opie*), 4309 (*Ji*), and 4312 (*Ji*). Maize genotype numbers (H1–H16) reported in Supporting Information 1. M—Thermo Scientific GeneRuler DNA Ladder Mix (100–10,000 bp)

**Table 4** Number of bands, monomorphic, polymorphic bands and percentage of polymorphism detected in 16 maize hybrids

Primer ID	RTE	Total Number of bands	Monomorphic	Polymorphic	Polymorphic bands (%)
4309	Ji	49	17	32	65
4310	Ji	30	2	28	93
4311	Ji	32	21	11	34
4312	Ji	38	27	11	29
		149	67	82	55
4303	Huck	56	31	25	45
4304	Huck	58	41	17	29
4306	Huck	44	11	33	7
4307	Huck	23	2	21	91
		181	85	96	53
4315	Opia	30	18	12	40
4317	Opia	45	22	23	51
4318	Opia	31	23	8	26
4319	Opia	20	2	18	90
		126	65	61	48
4320	Grande	29	8	21	72
4322	Grande	23	6	17	74
4323	Grande	21	4	17	81
4324	Grande	39	10	29	74
		112	28	84	75
4325	Tekay	19	3	16	84
4326	Tekay	24	15	9	37
4327	Tekay	39	13	26	67
4328	Tekay	27	9	18	67
		109	40	69	63
Total		677	285	392	
Average		34	23	20	58

### IRAP markers based on the *Opie* family

Seven LTR primers for the *Opie* RLX were tested in maize hybrids, of which primers 4315–4319 were suitable for scoring amplicons. A total of 126 bands were evaluated, of which 61 were PBs with a mean polymorphism of 48% (Table 4). The *Opie* LTR primers detected five positive unique bands with molecular sizes of 560, 720, 730, 1,031, and 2050 bp for hybrids H-16, H-7, H-14, H-16, and H-13, respectively. LTR primer 4317 detected one unique negative band of 720 bp for hybrid H-9, whereas primer 4319 distinguished H-13 with five unique negative bands with molecular sizes of 1080, 1090, 1400, 1510, and 1800 bp.

### IRAP markers based on the *Grande* element

Five LTR primers for the *Grande* RLX were tested in the maize hybrids, four of which (4320–4324) enabled molecular genetic evaluation. In all, 112 bands were detected, using *Grande* LTR primers, of which 84 were PBs with a mean polymorphism of 75%; the bands ranged from 250 to

3100 bp (Table 4). Six unique positive bands with molecular sizes of 705, 1200, (1400, 650), (1020, 1650), and 1031 bp characterized H-14, H-7, H-16, H-7, and H-8, respectively. In contrast, six unique negative bands characterized H-15, H-11, H-10, and H-16 with molecular sizes of 490 (610, 805), 250, and 795 bp, respectively (Table 5).

### IRAP markers based on the *Tekay* element

Six LTR primers for the *Tekay* RLX were tested in the 16 hybrids, four of which gave rise to suitable amplification for scoring (4325–4328). A total of 109 bands were generated from the four primers against the 16 maize hybrids, and ranged from 400 to 4100 bp. The number of PBs was 69, with a mean polymorphism percentage of 63% (Table 4). Three positive unique bands of 2900, 560, and 750 bp characterized H-5, H-16, and H-6, respectively, while four unique negative bands were detected with molecular sizes of 1040, 1590, and (1180, 2400) that characterized H-5, H-16, and H-1, respectively (Table 5).

**Table 5** Positive and negative unique bands as revealed by IRAP marker system

Primers For each element	Elements % of polymorphism for	Total	Unique bands				
			Tagged hybrids	Negative	Positive bands		
Band (bp) hybrids	Tagged						
4309	Ji	2	680	H10	298	H11	55.03
4310	Ji	2	2550	H7	–		
			1120	H16			
4312	Ji	2	410, 500	H16	–		
4303	Huck	2	1031	H15	2400	H5	53.03
4304	Huck	1	1600	H11			
4315	Opie	2	560	H16	–	48.41	
			720	H7			
4317	Opie	2	730	H14	720	H9	
4319	Opie	7	1031	H16	–		
			2050	H13	1080		
1090							
1400							
1510							
1800							
H13							
4320	Grande	1	705	H14	–	75.0	
4322	Grande	4	1200	H7			
490							
H15							
			1400, 650	H16			
4323	Grande	2		610, 805			
H11							
4324	Grande	6	1020, 1650	H7	250	H10	
					805	H11	
			1031	H8	795	H16	
4325	Tekay	1	2900	H5	–	63.30	
4326	Tekay	1		1040	H5		
4327	Tekay	3	560	H16			
			750	H6	1590	H16	
4328	Tekay	2		1180			
2400							
H1							
Total	20	40	22	18			

### Identifying maize hybrids with unique IRAP markers

Altogether (Table 5) 40 unique IRAP markers were generated from the 20 primers used, including 22 positive and 18 negative unique bands. Primers 4319 and 4324 produced the highest number of unique bands (7 and 6 bands, respectively). In contrast, the *Grande* RLX showed the highest number of unique bands (13), while the *Huck* RLX showed the smallest number of bands (3). The *Ji*, *Opie*, and *Tekay* RLXs showed 6, 11, and 7 unique bands, respectively. Table 5 shows that primer 4324 clearly separated

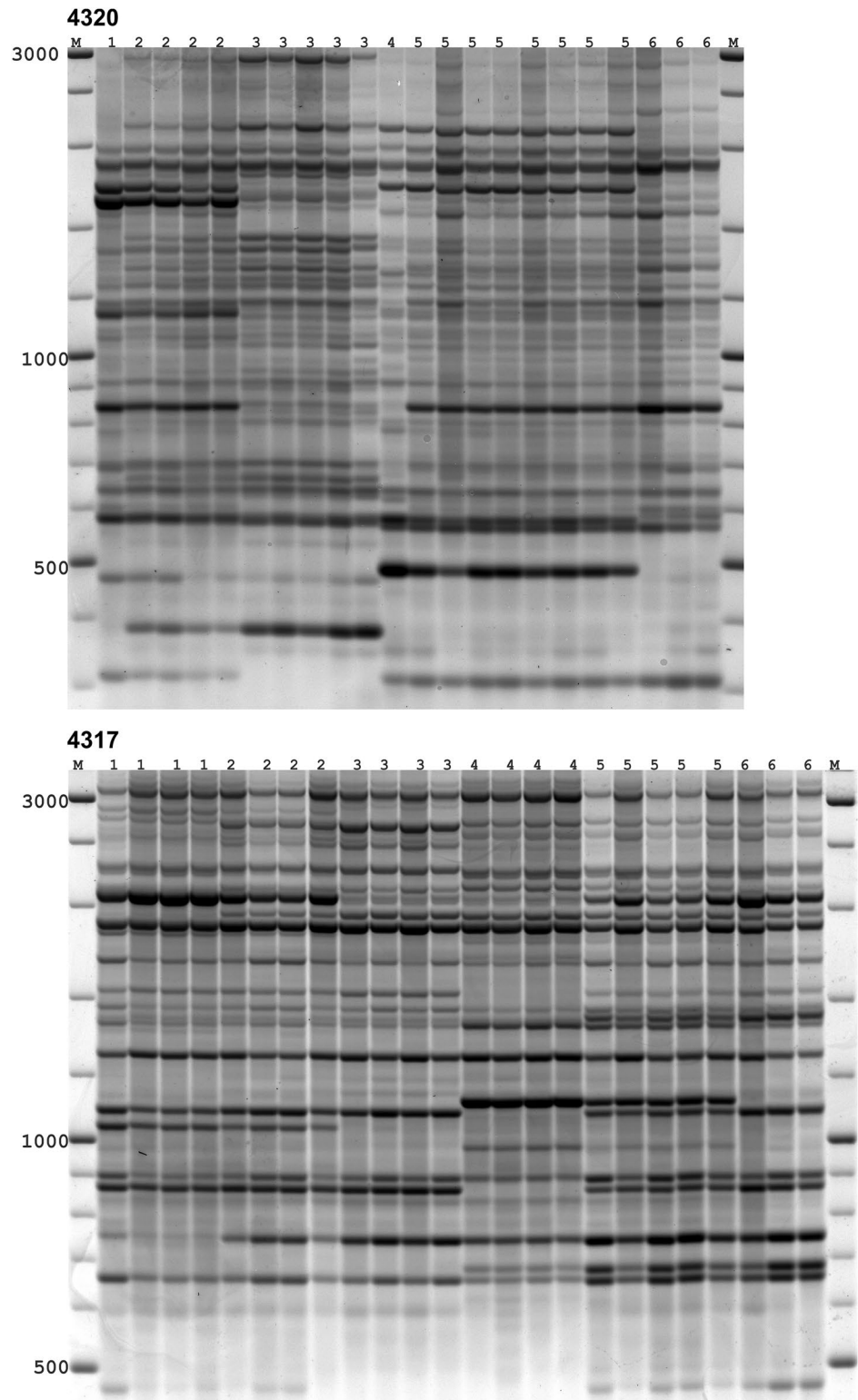
five hybrid lines and LTR primer 4322 distinguished three hybrids, while LTR primers 4309, 4310, 4303, 4315, 4317, and 4327 distinguished only two hybrid lines. LTR primers 4312, 4304, 4319, 4320, 4323, 4325, 4326, and 4328 characterized one hybrid each, while primers 4311, 4306, 4307, and 4318 could not separate any lines (Table 5).

The IRAP analysis used in this study succeeded in demonstrating positive and negative unique markers that aided in genotype discrimination. In all, 16 out of the 20 primers used revealed 40 unique IRAP markers, of which four (4311, 4306, 4307, 4318) did not produce any unique bands. The

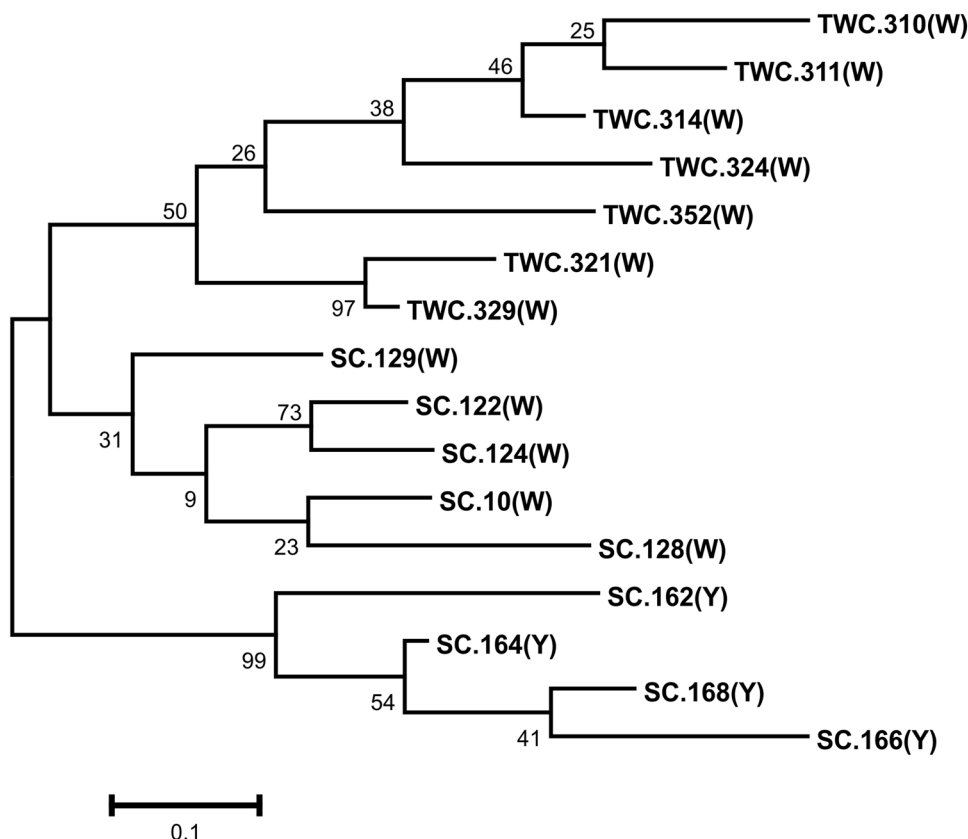
results revealed that some IRAP bands represented markers that are restricted to some hybrids. No unique bands were detected in hybrids H-2, H-3, H-4, and H-12, while other hybrid lines (H-7 and H-16) were identified by several primers. These unique IRAP bands could have several potential

uses for the study of maize genetic resources and for determining intellectual property rights for maize varieties.

**Fig. 2** Agarose gel electrophoresis patterns of IRAP amplicons obtained with primers 4317 (Opie) and 4320 (Grande) for individual seeds maize parental inbred lines (A619, A632, B73, Mo17) and their hybrids (A619 x A632 and B73 x Mo17). Maize parental inbred lines and their hybrids: 1—A632 (PI587140); 2—hybrid A619 x A632 (Ames23710); 3—A619 (PI587139); 4—Mo17 (PI558532); 5—hybrid B73 x Mo17 (Ames19097); 6—B73 (PI550473). M—Thermo Scientific GeneRuler DNA Ladder Mix (100–10,000 bp)



**Fig. 3** Dendrogram of maize genotypes generated by the IRAP primers



### Genetic relationships among maize hybrids

Understanding the relationships among genotypes within and between species has valuable applications in crop improvement programs. For this task, we selected two standard maize hybrids (A619 x A632 and B73 x Mo17) and their parental inbred lines (A619, A632, B73, Mo17) to evaluate the effectiveness of the IRAP method. Single-seed-derived DNA samples exclusively were used in this analysis. The bands shared between a hybrid and its parent inbred lines (Fig. 2) are clearly visible and also reflect by their decreased brightness the allelic dosage in the hybrid compared with the parent. For example, primer 4317 (Opie) yielded about 45 bands that could be detected well for all these maize lines, of which 25 were polymorphic (45%). Similar for primer 4320 (Grande), about 32 bands were well detected for all these maize lines, of which 16 were polymorphic (50%).

The IRAP banding profiles, which displayed from 21 to 95% polymorphism, were used to identify genetic similarity in the tested maize hybrid lines (Table 5). The highest similarity value (95%) was observed between the two white triple-cross hybrids (H-11 and H-12), which possess a common ancestor and share two parents, as seen in their pedigree (Table 1). In contrast, the lowest genetic similarity value (21%) was detected between the single-cross hybrid (H-4)

and the triple-cross white maize (H-11), indicating that these two hybrids were the most divergent genotypes. This dissimilarity can be attributed to the two genotypes inheriting their genetic makeup from different ancestors (Table 1). Genotypes that have low genetic similarity are of great interest for maize breeders. Weising et al. [65] mentioned that it is mandatory that genetically divergent parents be chosen that exhibit sufficient numbers of polymorphisms, but are not so distant as to cause sterility of the progeny. Estimation of genetic similarity based on molecular data is dependent on several factors, such as the number of markers analyzed, their distribution throughout the genome, and the quality of marker scoring. It is difficult to compare genetic distance between different studies, due to the difference in materials, number of genotypes analyzed, the number of alleles detected per marker, and the genetic diversity of the markers.

The genetic similarity coefficient determined for the maize hybrids was employed to develop a dendrogram based on IRAP data, as shown in Fig. 3. Cluster analysis resolved the 16 maize hybrids into two main clusters (A and B). The first cluster (A) was divided into two subclusters (C and D). The first subcluster (C) contained hybrid H-6 as a separate group, while the other subcluster (D) contained two groups; the first of which included H-4 and H-5, while the other contained H-9 (white single-cross) as a separate group. All yellow single-cross maize hybrids were clearly grouped in

the main cluster A. In contrast, cluster B was divided into two subclusters (E and F). Subcluster E contained all white triple-cross maize hybrids (H-11, H-12, H-10, H-13, H-14, and H-15), whereas the yellow triple-cross (H-16) was separated into a different group. Subcluster F contained all white single-cross maize hybrids in a separate group, while H-7 branched out into a different group.

## Conclusion

Here, we developed and applied a high-throughput IRAP technique for five LTR retrotransposon families to detect genetic polymorphisms among maize germplasm. These RLX families included *Opie*, *Ji*, *Cinful*, *Huck*, and *Grande*, which together comprise a large fraction of the maize genome. The RLX polymorphism captures the record of integration events, which are driven by retrotransposon activation and replication, that have been fixed in the germ line and inherited, and their subsequent fate in plant populations. The main goal of this study was to find efficient and high-throughput LTR retrotransposon markers for diversity analyses and to assess the polymorphism of these markers among maize genotypes originating from differing ecogeographical origins.

We successfully characterized the maize genotypes in worldwide and Egyptian collections, using high-throughput IRAP fingerprinting DNA markers. The DNA analysis of the lines of maize germplasm showed that even single LTR primers can be successfully used in the assessment of genetic differences at the line level and display several advantages, such as robustness, informativeness, and efficiency in breeding selection. We demonstrated here that the IRAP marker system provides a useful and simple electrophoretic technique for studying genetic diversity in maize, as they have in other plant species. The LTR primers used yielded multilocus fingerprints, displaying sufficiently high levels of polymorphism to differentiate between maize accessions and grouping them according to their cross level and kernel color. The markers are informative, reliable, and inexpensive for maize breeders and researchers. The number of differences between maize lines was sufficient to easily identify them as separate genotypes, correlated with their phenotypic differences.

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## Compliance with ethical standards

**Conflict of interest** The authors declare that they have no conflict of interest. All authors read and approved the final manuscript.

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